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Polymorphism of three milk protein genes in Mexican Jersey cattle

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1. Introduction

Mexican dairy cattle populations have been developed from imported genetic resources worldwide, incorporating a wide base of milk protein genes, including αS1-casein, αS2-casein, β-casein, κ-casein, α-lactalbumin and β-lactoglobulin. Particularly, the Jersey cattle populations have been constituted by American, Canadian [1], European, Australian and New Zealander progenitors. Jersey populations of these countries have been reported with variable gene frequencies for milk protein genes [2,3,4].

The importance to detect the genetic polymorphism for milk protein genes in dairy cattle populations is their association with cheese yield, rennet time, and curd firmness [3,5,6]. Besides, most of the available technologies have breed specific developments; therefore, when this technology is applied on breeds different than the one used as a model, imprecisions at the presence of genetic marker level from 8.3 to 54.9% might appear [7,8]. Molecular technologies have been developed to detect alleles and frequencies within protein milk genes, including specific PCR sequences, restriction enzymes, and actually single nucleotide polymorphism [9,10,11]. The development of breed specific SNPs is necessary for genotyping and association mapping to milk traits. The objectives were to determine the allelic and genotypic frequencies, genetic diversity and polymorphic information for β-casein, κ-casein, and β-lactoglobulin in Mexican Jersey cattle populations.

2. Materials and methods

2.1. Samples for DNA extraction

Samples were collected from 453 Jersey individuals registered by the Mexican Jersey Cattle Association, originated from Canadian, U.S., New Zealand, Australian and Mexican progenitors, including 401 cows and 52 sires. Sampled cows had at least a calf, whereas sires needed to have at least two calves in different herds. DNA samples from cows were obtained from blood and frozen semen was used to obtain the DNA in sires.

2.2. Selection of SNP primers

Primers used to genotype individuals were designed using OligoAnalyzer 3.1® (Integrated DNA Technologies, 2012, Iowa, USA), corresponding to the exons that represent the open reading frame of β-casein (β-CSN-encoding gene CSN2), κ-casein (κ-CSN-encoding gene CSN3), and β-lactoglobulin (β-LGB-encoding gene LGB) milk proteins reported in GenBank {NC_007304.4, http://www.ncbi.nlm.nih.gov/genbank; [12]}. The changing nucleotides were marked with a different fluorophore at the SNP position to distinguish each one during the allele identification by real time PCR. The reverse sequences and their complements, the coefficients of hairpin formation, autodimerization and creation of heterodimers, the
percentage of each nucleotide, and the fusion point for each sequence, necessary to design the thermo-cycling protocol, were determined with the same software.

Total SNP primers for whole genes were 28, including 8 primer sequences for β-lactoglobulin, 10 for κ-casein and 10 for β-casein. Primers were synthesized by KBioscience (Massachusetts, USA).

Molecular validation was done by amplifying the previously designed primers to corroborate the in silico performance. With the SNP primers that amplified well, the resolution curve was verified to guarantee the correct measure of the sample.

2.3. Genotyping of DNA

The PCR amplification was done with 4.11 µL of reaction solution (KASPar V4.0 Master Mix, KBioscience, USA) and 20 ng of genomic DNA; for the negative template control, 4 µL of molecular grade water was used. The polymorphisms were identified with the UV–Vis of the rtPCR thermo cycler (CFX 96, Bio-Rad, California, USA). Different fluorescence of SNP primers were used to detect the changing nucleotides. The thermocycling protocol was: one cycle at 94°C for 15 min, 10 cycles at 94°C for 20 s and 65°C for a min, with a decrease of 0.8°C per cycle in the second step, and 35 cycles at 94°C for 20 s, followed by a cycle at 57°C for 20 s, including the fluorescence report for each cycle in the last step. With the 28 validated SNPs, it was possible to identify the most common alleles for the three milk proteins. This increased the probability of detection of those alleles previously reported as being of low frequency in Bos taurus populations [13].

2.4. Data analysis

Polymorphic amplicons were considered to estimate the allelic diversity and effective number of alleles. Allelic and genotypic frequencies were estimated using the software Popgen32 [14].

3. Results and discussion

3.1. Genotyping for the three milk proteins

Data of haplotype and genotype frequencies of three protein milk genes in 453 individuals representing the Mexican Jersey cattle gene pool, are reported.

3.1.1. β-Casein

The alleles identified were A1, A2, A2, B and C (Table 1); A2 was the most frequent in the Mexican Jersey populations, 0.71. The frequency for this allele was similar to those reported, 0.58 to 0.65, for other Jersey populations [15,16,17]. This result suggests an absence of genetic selection for β-casein in Jersey cattle, including the Mexican populations. Some researchers have shown that the presence of the A2 allele in dairy cattle produces high quality milk associated with diminished cholesterol and triglycerides in humans [18,19]. The fact that A2 allele and A2A2 genotype were high in the Mexican Jersey populations is an important and distinctive aspect of this breed that could be used to improve the margin of profit for the milk producers. The frequency for the A2 allele in this study, 0.19, was similar to that reported by Van Eenennaam and Medrano [2] in a US Brown Swiss population, 0.18, which also is close to the frequencies for this allele found in Guernsey and Jersey, breeds with high total solids in milk [16,20]. This is in contrast to the frequencies of the A1 allele reported for Holstein populations, 0.49 to 0.95 [21,22].

The frequency of the A2 allele was 0.05, similar to that reported by Ng-Kwai-Hang and Grosclaude [23], who found a frequency for this allele in US Holstein-Friesian and Brown Swiss of 0.04. Meanwhile, the B allele had low frequencies in the Jersey populations, 0.04, similar to the frequencies reported for Jersey in some American and German populations, as well as for American Brown Swiss [2,20,24]. The low allele frequencies of A2 and B in this study, and the results reported by different authors in dairy breeds, suggest that these polymorphisms have little importance for breed differences in dairy cattle.

The most common genotype for β-casein was A2A2, 0.53. These results differ from those reported by Çardak [25] in a Turkish Holstein-Friesian population where the most frequent genotype was A1A1, 0.46. Furthermore, in this study the genotypes A1A2, A1B, and A1C were found, with frequencies lower than 0.05, which are similar to those reported by different authors, from 0.01 to 0.06, in US, New Zealand and Denmark Holstein, Brown Swiss and Ayrshire populations [16,20]. The genotypic frequencies found in the whole population were similar to those estimated only in females (Table 1). However, those corresponding to the male sub-population were different, with some absent genotypes (A1A1, A1B, A1C, A2A2, and BB). Hanusová et al. [26] found in Polish Holstein that one of the genotypes did not occur in the male subpopulation, even if it was present in the whole population. The absence of some alleles and genotypes in the male sub-population of this study might be a consequence of the reduced number of sampled individuals (n = 51 sires), even if the origin of sires is diverse.

3.1.2. κ-Casein

The alleles detected were A, B, and E (Table 2); B was the most common, 0.69. These findings were similar to the results reported for Colombian, German, and Chinese Jersey populations, 0.71 to 0.89 [11,27,28]. Some authors have shown that the presence of B allele in dairy cattle improves yield and quality of milk, raises milk casein fraction and diminishes whey protein fraction [3,6]. These results suggest an indirect genetic selection for the B allele in Jersey cattle, including the population studied, because the κ-casein alleles related to low total solid production, A (0.26) and E (0.05), are in lower frequencies than B allele. The frequency for the A allele in this population studied was similar to those reported by Trujillo et al. [27] and Ren et al. [11] in Jersey populations in Colombia and China, 0.26 in both populations, as well as in studies with Normande and Guernsey populations [27]. On the contrary, in Holstein there have been reported high frequencies, 0.68 to 0.89, for the A allele in several populations [22,29].

Jann et al. [28] and Boetcher et al. [30] reported in Holstein population frequencies for the E allele of 0.08 and 0.32, respectively. The frequencies for the A and E alleles in the Mexican Jersey populations, could be explained by the objective of selection over time for this breed, total solid production, common among most of the Mexican Jersey based milk producers. This is because these alleles influence positively milk production and negatively total solids in milk, and were indirectly selected against the genetic pool in Jersey.

Table 1

Allelic and genotypic frequencies of β-casein genes in the Mexican Jersey cattle populations.

<table>
<thead>
<tr>
<th>n</th>
<th>Allele frequencies</th>
<th>Genotype frequencies</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cows</td>
<td>401</td>
<td>0.22</td>
</tr>
<tr>
<td>Sires</td>
<td>52</td>
<td>0.12</td>
</tr>
<tr>
<td>Total</td>
<td>453</td>
<td>0.19</td>
</tr>
</tbody>
</table>

The most common genotype frequency for the κ-casein in this study was BB, 0.45. This result differs from those reported by Hernández et al. [31] and Cervantes et al. [32] in Holstein and Criollo Lecheró Tropical Mexican populations, where the most frequent genotype was AA, 0.57 and 0.62. Additionally, in the present study the genotypes BE and EE were found, with frequencies lower than 0.1, which were similar to the 0.08 reported by Boetcher et al. [30] for an Italian Holstein population.

### 3.1.3. β-Lactoglobulin

The alleles found in the sampled cows and sires were A, B, and C (Table 3). Allele B had the highest frequency, 0.72, higher than the estimates reported by Micifski et al. [17], Meza-Nieto et al. [33], and Ren et al. [11] in Holstein, Braunvieh, Jersey, and Criollo Lecheró Tropical populations, 0.31 to 0.65. Since the B allele of β-lactoglobulin is related to a high production of total solids, and given its relatively scarce and recent information on the polymorphism for this locus, the differences among the frequencies for this allele in several populations worldwide, suggest selection against it, by the means of selection in favor of the B allele of κ-casein gene in dairy cattle. The frequency observed for the A allele, 0.26, was similar to that reported by Ren et al. [11] in a Chinese Jersey population.

The frequency estimated for the C allele, 0.02, was similar to 0.01 reported by Berry et al. [4] in a New Zealand herd of F1 Holstein x Jersey. The result from the present study could be explained because some of the sampled cows in the Mexican Jerse population were imported from New Zealand.

The most frequent genotype for β-lactoglobulin in the present study was BB, 0.54. This result is similar to the findings of Meza-Nieto et al. [33] in Criollo Lecheró Tropical, where the frequency for the BB genotype was 0.53. In Turkish and Mexican Holstein populations, several authors have found frequencies between 0.24 and 0.38 [22,29,33].

Both genotypes BC and AA had a relatively low frequencies, 0.02 and 0.09, which were similar to 0.01 and 0.16 reported by Berry et al. [4] for a New Zealand F1 Holstein x Jersey population used to produce milk with high percentage of total solids.

### 3.2. Genetic diversity

The analyzed loci were 100% polymorphic, similar to the findings for Criollo Colombiano, Brahman, Holstein, Braunvieh, and Criollo Lecheró Tropical populations [32]. These results suggest that the loci analyzed remain without change regardless of breed or origin of the population.

The most polymorphic gene was β-casein with five alleles. The following more polymorphic genes were κ-casein and β-lactoglobulin.

### Table 2

<table>
<thead>
<tr>
<th>n</th>
<th>Allele frequencies</th>
<th>Genotype frequencies</th>
</tr>
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<tbody>
<tr>
<td></td>
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</tr>
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</tr>
<tr>
<td>Total</td>
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The most common genotype frequency for the β-lactoglobulin in the present study was BB, 0.45. This result differs from those reported by the estimates of Meza-Nieto et al. [17], Ren et al. [11] in Holstein, Braunvieh, Jersey, and Criollo Lecheró Tropical populations, 0.31 to 0.65. Since the B allele of β-lactoglobulin is related to a high production of total solids, and given its relatively scarce and recent information on the polymorphism for this locus, the differences among the frequencies for this allele in several populations worldwide, suggest selection against it, by the means of selection in favor of the B allele of κ-casein gene in dairy cattle. The frequency observed for the A allele, 0.26, was similar to that reported by Ren et al. [11] in a Chinese Jersey population.

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Both genotypes BC and AA had a relatively low frequencies, 0.02 and 0.09, which were similar to 0.01 and 0.16 reported by Berry et al. [4] for a New Zealand F1 Holstein x Jersey population used to produce milk with high percentage of total solids.

### 3.3. Heterozygosity and Hardy–Weinberg equilibrium

The studied population showed up differences (P < 0.05) between observed (Hₒ) and expected (Hₑ) heterozygosity (Table 4). These findings indicate that Hₒ doubled Hₑ in populations where reproductive management and genetic improvement programs have been applied. Similar results were reported by other authors in Holstein, Brahman, Braunvieh and Jersey populations [11,31]. In those populations matings were planned, therefore some influence from artificial selection for an economic trait could be expected.

### 4. Conclusion

The sampled Mexican Jersey population has a diverse genetic pool. The most common alleles were A² for β-casein, and B for both κ-casein and β-lactoglobulin. The Mexican Jersey population has the potential to be used in genetic improvement programs, aimed to improve the milk quality traits of economic importance.

### Financial support

This project was supported by CONAGEN (2013-001) (Mexican Council of Livestock Genetic Resources) and Universidad Autónoma Chapingo, Mexico (DGIP-11550301). The authors also thank CONACYT for the financial support to the first author during his Master of Science studies.

### References


### Table 3

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<td>453</td>
<td>0.26</td>
</tr>
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### Table 4

<table>
<thead>
<tr>
<th>Locus</th>
<th>Hₒ</th>
<th>Hₑ</th>
<th>X²</th>
</tr>
</thead>
<tbody>
<tr>
<td>CSN2</td>
<td>0.60</td>
<td>0.46</td>
<td>0.033*</td>
</tr>
<tr>
<td>CSN3</td>
<td>0.50</td>
<td>0.45</td>
<td></td>
</tr>
<tr>
<td>LGB</td>
<td>0.50</td>
<td>0.41</td>
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with three observed alleles each one. The three genes had one effective allele. This is, when comparison for the average observed alleles to the average effective alleles through the three loci was done, the number of effective alleles was 30% less than those observed.

* X² test P value (P value < 0.05, not consistent with Hardy Weinberg Equilibrium).


